

A Kinetic Model of Molecular Titration

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1 Motivation

One of the components of our Circuit Control Toolbox is a suite of parts which allow for the use of decoy transcription factor binding arrays that make use of Molecular Titration to shift the sensitivity of a gene or circuit's response to transcription factor concentration. Part of our suite is a series of parts which can be used to construct tetO and lacO binding arrays of arbitrary length using our protocol for UNS-guided Iterative Capped Assembly.

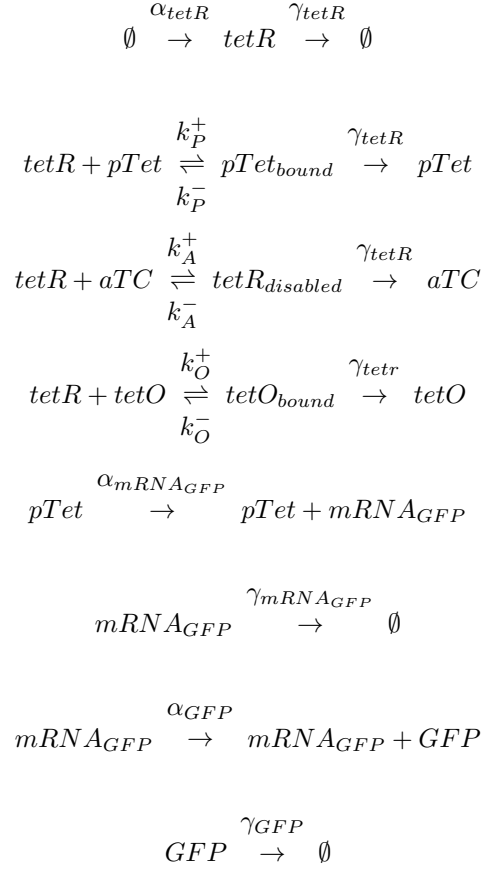
For teams and scientists interested in designing their own decoy binding arrays, it is important to be able to make informed predictions about the part's impact on an existing circuit prior to its actual construction. To facilitate this, we developed a mathematical model which is explicitly parametrized by physiological values for variable aspects of Toolbox parts, such as the number of binding sites on an array or the array's plasmid backbone.

2 Model

2.1 Design and Solution

We developed a simple kinetic ODE model of a basic GFP-expression system, where constitutively expressed tetR inhibits the expression of a pTet-regulated GFP. The model accounts for the activity of aTC, which disables tetR's DNA-binding ability when bound, and the presence of

decoy tetO arrays. The kinetic diagram for our model is given by:



which lead to the mass-action ODE system

$$\begin{aligned}
\dot{T} &= \alpha_T - \gamma(T + TP + TA + TO)T - \\
&\quad k_P^+ T * P + k_P^- TP - k_A^+ T * A + k_A^- TA - k_O^+ T * O + k_O^- TO \\
\dot{P} &= -k_P^+ T * P + k_P^- TP \\
\dot{TP} &= k_P^+ T * P - k_P^- TP \\
\dot{A} &= -k_A^+ T * A + k_A^- TA \\
\dot{TA} &= k_A^+ T * A - k_A^- TA \\
\dot{O} &= -k_O^+ T * O + k_O^- TO \\
\dot{TO} &= k_O^+ T * O - k_O^- TO \\
\dot{m} &= \alpha_m P - \gamma_m m \\
\dot{G} &= \alpha_G m - \gamma_G G
\end{aligned}$$

where for simplicity we have written $T \equiv tetR$, $P \equiv pTet$, $TP \equiv pTet_{bound}$, $A \equiv aTC$, $TA \equiv tetR_{disabled}$, $O \equiv tetO$, $TO \equiv tetO_{bound}$, $m \equiv mRNA_{GFP}$, and $G \equiv GFP$.

Note that we can naturally define conserved quantities in this system which will be invariant over time. These are

$$\begin{aligned}P_{tot} &= P + TP \\A_{tot} &= A + TA \\O_{tot} &= O + TO\end{aligned}$$

Furthermore, note that one can deduce from the equations that we can intuitively define $T_{tot} = T + TP + TA + TO$. This allows us to define the flux

$$\begin{aligned}\dot{T}_{tot} &= \dot{T} + \dot{TP} + \dot{TA} + \dot{TO} \\ \dot{T}_{tot} &= \dot{T} - \dot{P} - \dot{A} - \dot{O} \\ \dot{T}_{tot} &= \alpha_T - \gamma_T T_{tot}\end{aligned}$$

where the second equality is true because for any conservation condition X_{tot} , $\dot{X}_{tot} = 0$.

We are now able to reduce our system of 9 variables down to 6 variables using our conservation conditions:

$$\begin{aligned}\dot{T}_{tot} &= \alpha_T - \gamma_T T_{tot} \\ \dot{P} &= -k_P^+ T * P + k_P^- (P_{tot} - P) \\ \dot{A} &= -k_A^+ T * A + k_A^- (A_{tot} - A) \\ \dot{O} &= -k_O^+ T * O + k_O^- (O_{tot} - O) \\ \dot{m} &= \alpha_m P - \gamma_m m \\ \dot{G} &= \alpha_G m - \gamma_G G\end{aligned}$$

This expression can be evaluated at steady-state by setting all of the time derivatives on the left hand side to 0 and rearranging terms to obtain

$$\begin{aligned}T_{tot;ss} &= \frac{\alpha_T}{\gamma_T} \equiv J_T \\ P_{ss} &= \frac{K_P P_{tot}}{K_P + T_{ss}} \\ A_{ss} &= \frac{K_A A_{tot}}{K_A + T_{ss}} \\ O_{ss} &= \frac{K_O O_{tot}}{K_O + O_{ss}} \\ m_{ss} &= \frac{\alpha_m}{\gamma_m} P_{ss} \equiv J_m P_{ss} \\ G_{ss} &= \frac{\alpha_G}{\gamma_G} m_{ss} \equiv J_G m_{ss}\end{aligned}$$

Where $K \equiv k^-/k^+$ is the dissociation constant, and we have defined flux terms J for convenience of notation.

In order to write an expression for G_{ss} in terms of only the rate parameters and conservation conditions of the system, we can see from the above equations that we need to determine an expression for T_{ss} in terms of only rate parameters and conservation conditions. We can do this using the T_{tot} expression: recall that we had defined

$$T_{tot} = T + TP + TA + TO = T + (P_{tot} - P) + (A_{tot} - A) + (O_{tot} - O).$$

This means that

$$T_{tot;ss} = J_T = T_{ss} + (P_{tot} - P_{ss}) + (A_{tot} - A_{ss}) + (O_{tot} - O_{ss}),$$

and we can substitute our expressions for P_{ss} , A_{ss} , and O_{ss} and rearrange terms to eventually obtain

$$0 = T_{ss}^4 + a_1 T_{ss}^3 + a_2 T_{ss}^2 + a_3 T_{ss} + a_4,$$

where

$$\begin{aligned} a_1 &= K_P + K_A + K_O - J_T \\ a_2 &= K_P K_A + K_P K_O - K_P J_T + K_A K_O - K_A J_T - K_O J_T + \\ &\quad K_P P_{tot} + K_A A_{tot} + K_O O_{tot} \\ a_3 &= K_P K_A K_O - K_P K_I J_T - K_P K_O J_T - K_A K_O J_T + \\ &\quad K_P P_{tot} (K_A + K_O) + K_A A_{tot} (K_P + K_O) + K_O O_{tot} (K_P + K_A) \\ a_4 &= K_P K_A K_O (P_{tot} + A_{tot} + O_{tot} - J_T). \end{aligned}$$

This provides an expression for which can be solved in a straightforward fashion for its maximal positive real root to compute the value of T_{ss} solely in terms of the rate parameters and conserved conditions of the system. Returning to our original reduced model, we see that once we obtain T_{ss} we can follow a cascade of function composition to G_{ss} via

$$G_{ss} = J_G m_{ss} = J_G (J_m P_{ss}) = J_G J_m \frac{K_P P_{tot}}{K_P + T_{ss}}.$$

2.2 Explicit Model Parametrization by Toolbox Parameters

We have now obtained an expression to compute the steady-state concentration of GFP in our system using only the values of the rate parameters and conservation conditions. Note that these conservation condition values correspond explicitly to physiological parameters of toolbox settings in the following way:

The two most easily-tunable parameters for the Titration suite of our Toolbox are the copy number of the plasmid backbone and the number of binding sites on the decoy binding array. We can define two parameters: B to define the mean copy number of the plasmid in a given cell, and C to define the number of copies on a single plasmid. Using the approximation that in an *E. coli* volume, 1 molecule \approx 1 nM concentration [1], we can decompose our conserved quantities as

$$\begin{aligned} P_{tot} &= C_{pTet} * B_{pTet} \\ O_{tot} &= C_{tetO} * B_{tetO} \end{aligned}$$

where we can have distinct values of C and B if the reporter and array are on different plasmids.

This decomposition provides our model with an explicit link between the design specifications of a decoy binding array with its kinetic realization. Details about the reported mean copy numbers for various BioBrick plasmid backbones are given in

http://parts.igem.org/Help:Plasmid_backbones/Nomenclature

2.3 Fluorescence to Concentration Conversion

Being a kinetic model, our model processes and evaluates the behavior of the system through concentrations of the relevant proteins. However, we and many users of the Toolbox would likely use fluorescent measurements rather than fluorescent protein concentrations to characterize the behavior of their system. In order to compare our model's results to experimental data, then, we would need a process to convert between fluorescence and fluorescent protein concentration.

Our objective is to obtain an absolute fluorescence unit called MEGFP (Molecules of Equivalent GFP). We modeled our procedure off the workflow developed by the Endy Lab [2] to convert arbitrary fluorescent measurements from a plate reader to MEGFP. We obtained a purified concentration of GFP [3] and diluted it down to various concentration levels and measured these values over different days on the plate reader. The relationship between GFP concentration and GFP fluorescence was linear, as previously described in the literature [4]. We obtained the conversion curve

$$Fluorescence_{MEGFP} = \frac{(Fluorescence_{PlateReader;au} + 683.02)}{100}$$

with an R^2 value of 0.93263. Here $Fluorescence_{MEGFP}$ takes units of concentration (nM).

We also wanted to be able to convert fluorescence values obtained via flow cytometry on our FACS machine to units of MEGFP. To do this, we measured an IPTG induction curve of a standard GFP-expression cassette driven by pLac in the presence of constitutive lacI expression (BBa K2066110) on the FACS machine and converted it to absolute fluorescence units (MEFL) using our standard FACS protocol [see Protocol section of our Wiki]. We then immediately measured these same samples in the plate reader and obtained fluorescence measurements there. We then plotted the plate reader fluorescence values against the FACS fluorescent values at each condition, and found that the the relationship was linear as expected [5]. We obtained the conversion curve

$$Fluorescence_{PlateReader;au} = 0.9054 * Fluorescence_{FACS;MEFL} + 918.10$$

with an R^2 value of 0.9642. This curve can be composed with the above curve to obtain a conversion from MEFL values to MEGFP concentrations.

3 Results

3.1 Model Tuning

In order to validate our model we used it to predict the impact of an 85-repeat tetO array (BBa K2066550) on pSB1C3 backbone on the aTC-induction curve of a pTet-GFP reporter construct in the presence of constitutive tetR expression (BBa K2066053).

We first searched the literature for values for rate constants. We obtained the following values:

Parameter	Meaning	Value	Justification
α_T	Production rate of tetR	? s^{-1}	Fit to Data (see next page)
γ_T	Degradation rate of tetR	0.6207288 s^{-1}	Cell doubling time in M9 minimal media is 67 minutes [6].
K_P	Dissociation constant of tetR with the pTet Promoter	0.1 nM	[7]
K_A	Dissociation constant of aTC with the tetR molecule	0.36 nM	[7]
K_O	Dissociation constant of tetR with the tetO binding site	0.1 nM	[7]
α_m	Transcription rate from the pTet promoter	? nM/s	Fit to Data (see next page)
γ_m	Degradation rate of the GFP transcript	1/120 s^{-1}	Assumption that transcript half-life is 2 minutes [8].
α_G	Translation rate of GFP from transcript	? nM/s	Fit to Data (see next page)
γ_G	Degradation rate of GFP	0.6207288 s^{-1}	Cell doubling time in M9 minimal media is 67 minutes [6].

Table 1: Known Parameter Values for Kinetic Model

Since we were unable to obtain parameter values for the three α terms with which we could be confident, we decided to fit these parameters to an aTC-induction curve of BBa K2066110 without the decoy binding array. We iterated through a parameter range that spanned three orders of magnitude for each of these α terms and minimized the pointwise squared error between the model-generated steady-state induction curve and the experimentally observed steady-state induction curve. The model-generated curves were obtained by solving our model for G_{ss} with $P_{tot} = 1 * 200$ and A_{tot} given by the aTC induction point. We obtained a best-fit parameter set of $\alpha_T = 416$, $\alpha_m = 1/311$, and $\alpha_G = 172$.

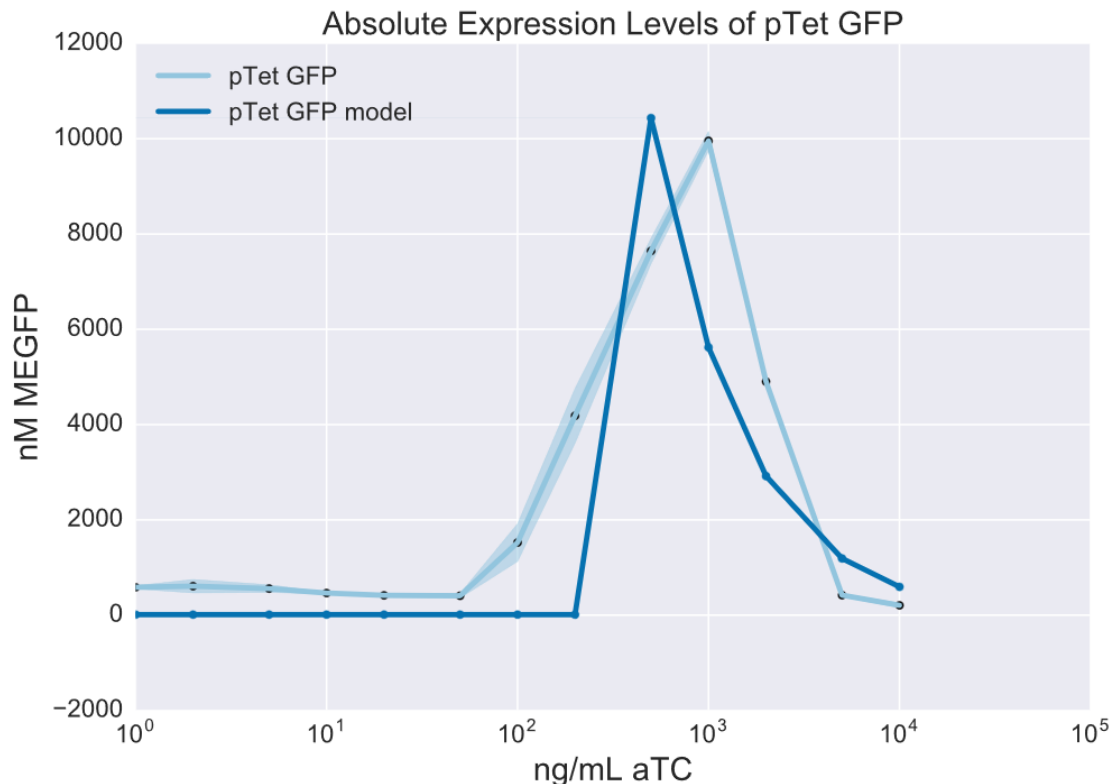


Figure 1: Optimized fit of Kinetic Model to observed induction curve of pTet GFP (BBa K2066053). $\alpha_T = 416$, $\alpha_m = 1/311$, and $\alpha_G = 172$. $P_{tot} = 1 * 200$ and $O_{tot} = 0$.

The model-generated induction curve’s rapid induction is a function of the constraints on the model’s dynamics imposed by the simplicity of its design, as this represents a sharp transition between two limiting regimes of behavior as the free tetR concentration crosses some threshold. Hence we felt that the fit could not be better-optimized on this dataset.

3.2 Validation of Predictive Power

We then fixed our optimized α parameters and included the 85x tetO array into our model. Because our decoy binding array has 85 repeats, and because pSB1C3 is reported to have 100-300 copies per cell [9], we follow our guidelines from Section 2.2 and set $O_{tot} = 85 * 200$. We obtained a good fit!

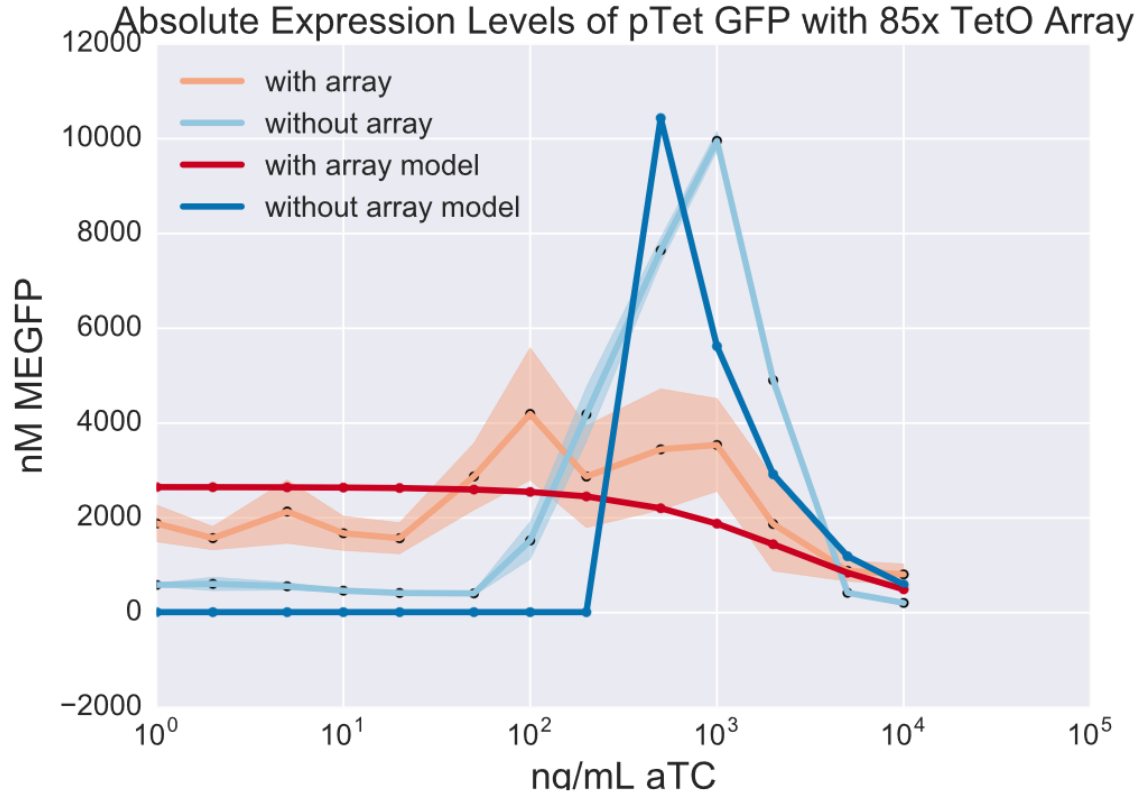


Figure 2: The Kinetic Model predicts the impact of the 85x tetO array on pSB1C3 on the induction curve of BBa K2066053 on pSB1A3. Shading on data curves represent mean \pm SEM.

4 Discussion

We have demonstrated that our simple kinetic model is able to capture the impact of the decoy binding array on the gene expression dynamics of a simple inducible reporter construct. We note that our model prediction suggests that the use of the 85x tetO array on pSB1C3 induces a sufficiently strong titration effect that it negates the impact of aTC induction, having the reporter construct express at its maximal level at the minimal aTC concentration. This is most likely caused by the constraints imposed by our model's design, namely its simplicity and its reliance on deterministic mass-action kinetics.

The 85x tetO binding array on pSB1C3 backbone specification likely lies on a graded transition between shifted, functional induction curves and shifted, nonfunctional induction curves imposed by excessive molecular titration, and hence the successful induction observed in our data is driven by the nuances of the stochasticity inherent in biological processes. Indeed if we predict the impact of a smaller binding array on a lower-copy plasmid, such as a 16x tetO array on pSB3K3 (Fig. 3), we observe that the predicted curve has shifted to the left but still induces. It will be interesting to see the predictions generated by stochastic models and to assess whether they match observed data to an even better extent.

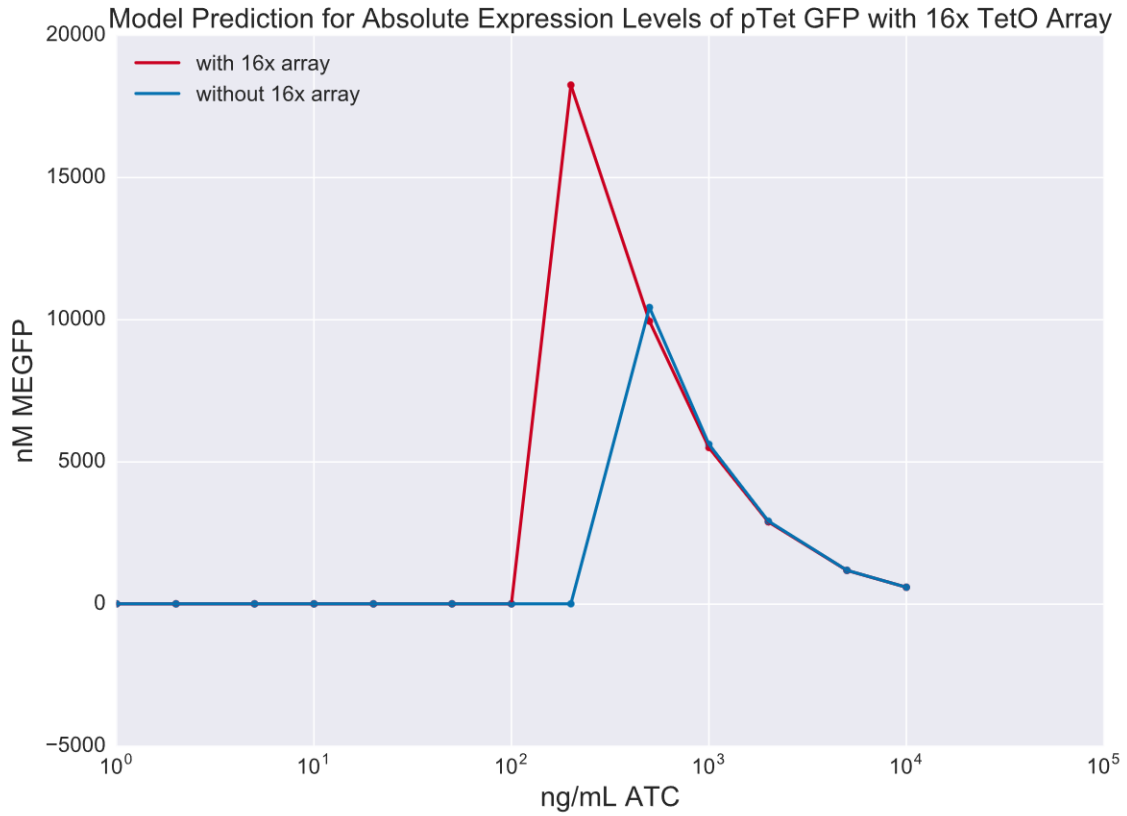


Figure 3: Model Predictions for the induction of BBa K2066053 on pSB1A3 backbone with or without 16x tetO array on pSB3K3. $P_{tot} = 1 * 200$ and $O_{tot} = 16 * 10$. α parameters are as before.

References

- [1] Milo, R., Phillips, R., & Orme, N. (2015). Cell biology by the numbers. Garland Science.
- [2] Kelly, J. R., Rubin, A. J., Davis, J. H., Ajo-Franklin, C. M., Cumbers, J., Czar, M. J., ... & Endy, D. (2009). Measuring the activity of BioBrick promoters using an in vivo reference standard. Journal of biological engineering, 3(1), 1.
- [3] <https://www.thermofisher.com/order/catalog/product/88899>
- [4] Furtado, A., & Henry, R. (2002). Measurement of green fluorescent protein concentration in single cells by image analysis. Analytical biochemistry, 310(1), 84-92.
- [5] Castillo-Hair, S. M., Sexton, J. T., Landry, B. P., Olson, E. J., Igoshin, O. A., & Tabor, J. J. (2016). FlowCal: A user-friendly, open source software tool for automatically converting flow cytometry data from arbitrary to calibrated units. ACS synthetic biology.
- [6] <https://www.genome.wisc.edu/resources/k12growth/mg1655growthcurve.htm>

- [7] Amit, R., Garcia, H. G., Phillips, R., & Fraser, S. E. (2011). Building enhancers from the ground up: a synthetic biology approach. *Cell*, 146(1), 105-118.
- [8] So LH, Ghosh A, Zong C, Sepúlveda LA, Segev R, Golding I. General properties of transcriptional time series in *Escherichia coli*. *Nat Genet*. 2011 Jun 43(6):554-60
- [9] <http://parts.igem.org/Part:pSB1C3>